

European Patent Office

Office européen des brevets

Bescheinigung

Certificate

Attestation

Die angehefteten Unterlagen stimmen mit der ursprünglich eingereichten Fassung der auf dem nächsten Blatt bezeichneten europäischen Patentanmeldung überein.

The attached documents are exact copies of the European patent application conformes à la version described on the following page, as originally filed.

Les documents fixés à cette attestation sont initialement déposée de la demande de brevet européen spécifiée à la page suivante.

Patentanmeldung Nr. Patent application No. Demande de brevet n°

97116863.8

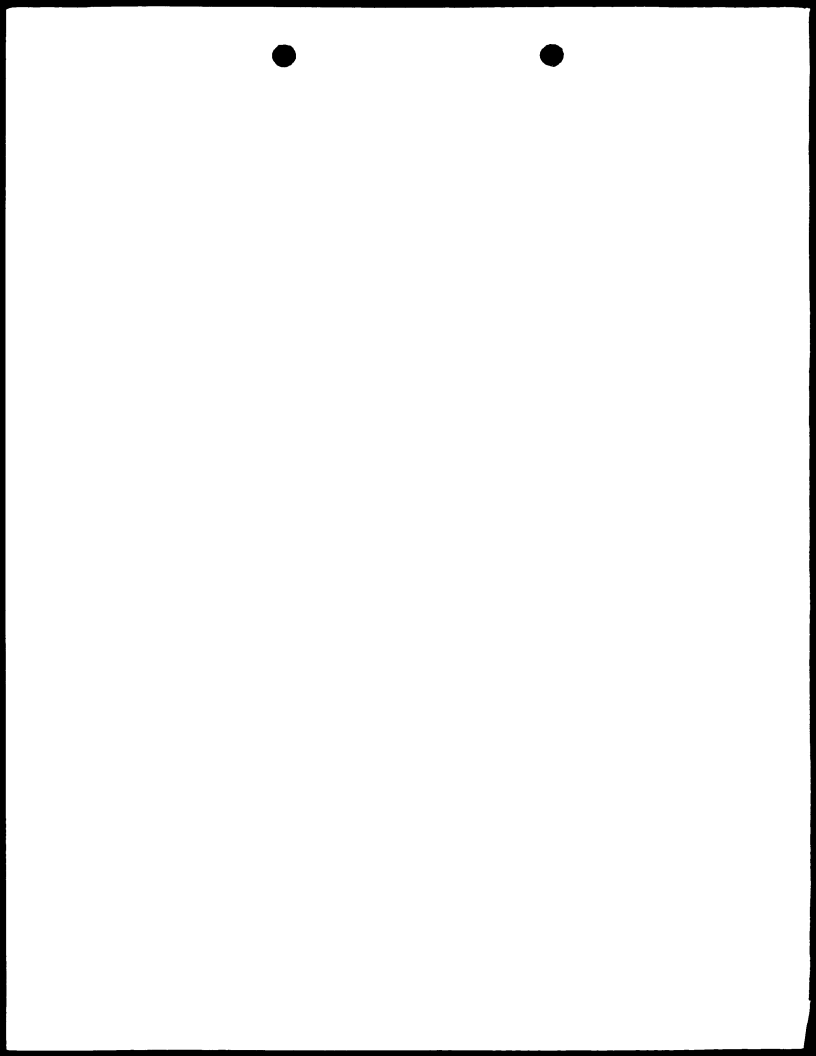
Der Prasident des Europäischen Patentamts: Im Auftrag

For the President of the European Patent Office

Le Président de l'Office européen des brevets

I.L.U. HATTEN-HECKMAN

DEN HAAG,DEN THE HASTER. LA HAYE, LE





Europäisches Patentamt **European Patent Office** 

Office européen des brevets

# Blatt 2 der Bescheinigung Sheet 2 of the certificate Page 2 de l'attestation

Anmeldung Nr Application no Demande n°

97116863.8

Anmeldetag Date of filing Date de dépôt

29/09/97

Anmeider
Applicant(s)
Demandeur(s)
APPLIED RESEARCH SYSTEMS ARS HOLDING N.V.
Curação

NETHERLANDS ANTILLES

Bezeichnung der Erfindung Title of the invention Titre de l'invention

Amino-terminal truncated c-c chemokines as chemokine antagonist

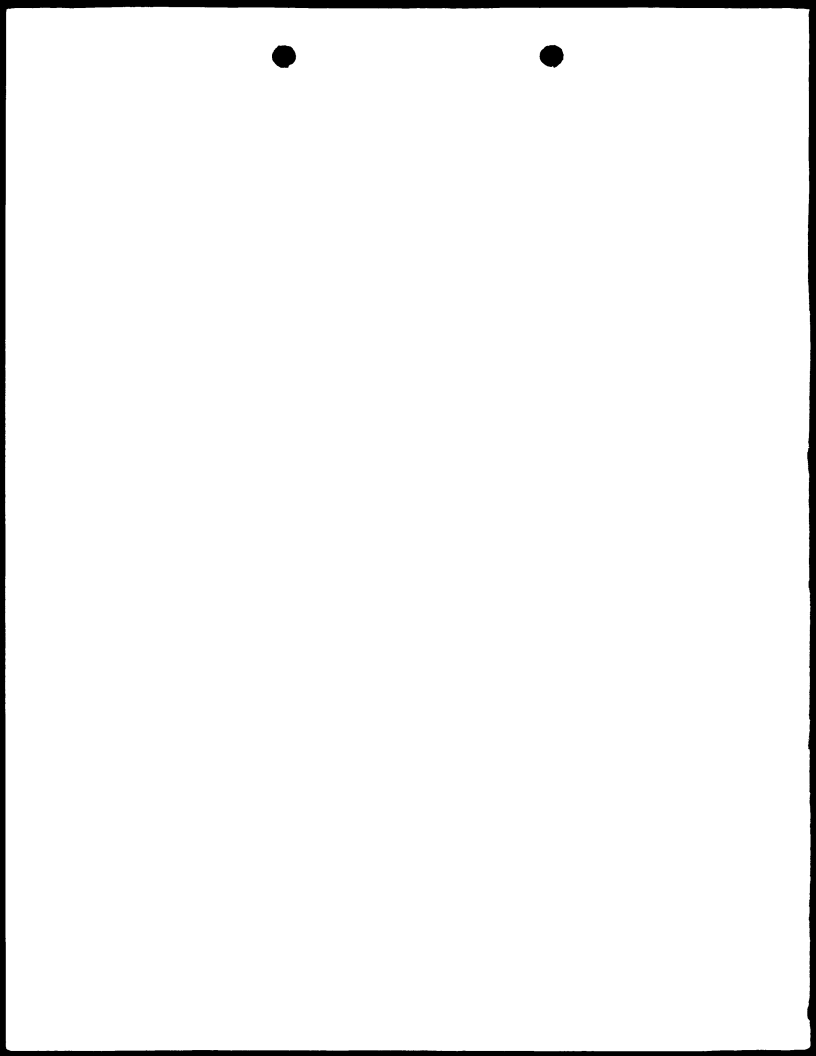
In Anspruch genommene Prioriat(en) / Priority(ies) claimed / Priorité(s) revendiquée(s)

Staat State Pays Tag Date Date Aktenzeichen File no Numéro de dépôt

Internationale Patentklassifikation International Patent classification Classification internationale des brevets

C12N15/19, C07K14/52, A61K38/19, A61K38/48, G01N33/68

 $\label{thm:local_problem} A m \ A n meldetag \ benan te \ Vertrag staten \\ Contracting \ states \ designated \ at \ date \ of \ filing \ A T/BE/CH/CY/DE/DK/ES/FI/FR/GB/GR/IE/IT/LI/LU/MC/NL/PT/SE/TR \\ A T/BE/CH/CY/DE/DK/ES/TR \\ A T/BE/CH/CY/DK/ES/TR \\ A T/BE/CH/CY/DK/ES/TR \\ A T/BE/CH/CY/D$ 



10

15

20

25



# AMINO-TERMINAL TRUNCATED C-C CHEMOKINES AS CHEMOKINE ANTAGONISTS

## FIELD OF THE INVENTION

The present invention relates to amino-terminal truncated C-C chemokines, having chemokine antagonistic activity. In particular, we refer to amino-terminal truncated C-C chemokines missing up to 5 amino-terminal amino acids and having a chemokine antagonistic activity. More particularly, MCP-2 (6-76) and RANTES (3-68) are described, as well as cDNA sequences encoding them, their use in therapy and/or in diagnosis of the diseases, in which an antagonistic activity of the chemokine effects is required, and pharmaceutical compositions comprising them.

### BACKGROUND OF THE INVENTION

Chemokines constitute a family of small pro-inflammatory cytokines with leukocyte chemotactic and activating properties. Depending on the position of the first cysteines, the chemokine family can be divided in C-C, C-X-C and C-X<sub>3</sub>-C chemokines (Baggiolini et al., Ann. Rev. Immunol., 55, 97-179, 1994; Baggiolini et al., Ann. Rev. Immunol., 15, 675-705, 1997; and Taub et al., Cytokine & Growth Factor Reviews, 7, 335-76, 1996).

Many C-X-C chemokines such as interleukin-8 (IL-8) are chemotactic for neutrophils, while C-C chemokines, such as monocyte chemotactic protein-3 (MCP-3), are active on a variety of leukocytes including monocytes, lymphocytes, eosinophils, basophils, NK cells and dendritic cells.

The NH<sub>2</sub> terminal domain of chemokines is involved in receptor-binding and NH<sub>2</sub>-terminal processing can either activate chemokines or render chemokines completely inactive.

The C-X-C chemokine platelet basic protein becomes a neutrophil chemotactic peptide (NAP-2) only after removal of the 24 NH<sub>2</sub>-terminal residues (Walz et al., *Biochem. Biophys. Res. Commun.*, 159, 969-75, 1989; and Van Damme et al., *Eur. J. Immunol.*, 20, 2113-8, 1990).

televiment in the control of the televiment of the control of

10

15

20

25

TTO OU ZOUDTTOOF#IL

-2-

SOO S FOODITODA

front of the first Cys in all neutrophil chemotactic C-X-C chemokines, causes complete inactivation (Clark-Lewis I. et al., J. Biol. Chem., 266, 23128-23134, 1991).

Similar NH<sub>2</sub>-terminal proteolysis (up to 8 amino acids) of another C-X-C chemokine, granulocyte chemotactic protein-2 (GCP-2), has no effect on the neutrophil chemotactic activity (Proost P.et al., *Biochemistry*, 32, 10170-10177, 1993).

The synthetical C-C chemokines MCP-1, MCP-3 and RANTES missing the 8 to 9 NH<sub>2</sub>-terminal amino acids are inactive on monocytes and are useful as receptor antagonists (Gong J. et al., *J. Biol. Chem.*, 271, 10521-7, 1996; and Gong G. et al., *J. Exp. Med.*, 631-40, 1995).

Extension of RANTES with one methionine results in complete inactivation of the molecule and Met-RANTES behaves as an antagonist for the authentic RANTES (Proudfoot A.E. et al., J. Biol. Chem., 271, 2599-2603, 1996).

#### DESCRIPTION OF THE INVENTION

The main object of the present invention are amino-terminal truncated C-C chemokines, having chemokine antagonistic activity. In particular, we refer to amino-terminal truncated C-C chemokines missing up to 5 amino-terminal amino acids and having a chemokine antagonistic activity. A non-limiting list of C-C chemokines include: MCP-1, MCP-2, MCP-3, MCP-4, MCP-5, RANTES, MDC, MIP-1 $\alpha$  and MIP-1 $\beta$ .

More particularly, one object of the present invention is MCP-2 (6-76), which is MCP-2 missing the 5 NH<sub>2</sub>-terminal amino acids, as shown in Figure 1.

Another object of the present invention is RANTES (3-68), which is RANTES missing the first 2 amino acids, as shown in Figure 1.

Such amino-terminal truncated C-C chemokines of the invention can be in a glycosylated or not-glycosylated form.

Another object of the invention are the DNA molecules comprising the DNA sequences coding for the amino-terminal truncated chemokines of the invention, including nucleotide sequences substantially the same.

"Nucleotide sequences substantially the same" includes all other nucleic acid sequences which, by virtue of the degeneracy of the genetic code, also code for the given amino acid sequences.

10

15

20

25

\_3-

The invention also includes expression vectors which comprise the above DNAs, host-cells transformed with such vectors and a process of preparation of such aminoterminal truncated chemokines of the invention, through the culture in appropriate culture media of said transformed cells.

The DNA sequence coding for the proteins of the invention can be inserted and ligated into a suitable plasmid. Once formed, the expression vector is introduced into a suitable host cell, which then expresses the vector(s) to yield the desired protein.

Expression of any of the recombinant proteins of the invention as mentioned herein can be effected in eukaryotic cells (e.g. yeasts, insect or mammalian cells) or prokaryotic cells, using the appropriate expression vectors. Any method known in the art can be employed

For example the DNA molecules coding for the proteins obtained by any of the above methods are inserted into appropriately constructed expression vectors by techniques well known in the art (see Sambrook et al, 1989). Double stranded cDNA is linked to plasmid vectors by homopolymeric tailing or by restriction linking involving the use of synthetic DNA linkers or blunt-ended ligation techniques: DNA ligases are used to ligate the DNA molecules and undesirable joining is avoided by treatment with alkaline phosphatase.

In order to be capable of expressing the desired protein, an expression vector should comprise also specific nucleotide sequences containing transcriptional and translational regulatory information—linked to the DNA coding the desired protein in such a way as to permit gene expression and production of the protein. First in order for the gene to be transcribed, it must be preceded by a promoter recognizable by RNA polymerase, to which the polymerase binds and thus initiates the transcription process. There are a variety of such promoters in use, which work with different efficiencies (strong and weak promoters).

For eukaryotic hosts, different transcriptional and translational regulatory sequences may be employed, depending on the nature of the host. They may be derived form viral sources, such as adenovirus, bovine papilloma virus. Simian virus or the like,

respression inxamples are the fix promoter of the rierpes whis the sixth carry

10

15

20

25

3()

-4-

promoter, the yeast gal4 gene promoter, etc. Transcriptional initiation regulatory signals may be selected which allow for repression and activation, so that expression of the genes can be modulated.

The DNA molecule comprising the nucleotide sequence coding for the protein of the invention is inserted into vector(s), having the operably linked transcriptional and translational regulatory signals, which is capable of integrating the desired gene sequences into the host cell.

The cells which have been stably transformed by the introduced DNA can be selected by also introducing one or more markers which allow for selection of host cells which contain the expression vector. The marker may also provide for phototrophy to a auxotropic host, biocide resistance, e.g. antibiotics, or heavy metals such as copper, or the like. The selectable marker gene can either be directly linked to the DNA gene sequences to be expressed, or introduced into the same cell by co-transfection. Additional elements may also be needed for optimal synthesis of proteins of the invention.

Factors of importance in selecting a particular plasmid or viral vector include: the ease with which recipient cells, that contain the vector may be recognized and selected from those recipient cells which do not contain the vector; the number of copies of the vector which are desired in a particular host; and whether it is desirable to be able to "shuttle" the vector between host cells of different species.

Once the vector(s) or DNA sequence containing the construct(s) has been prepared for expression the DNA construct(s) may be introduced into an appropriate host cell by any of a variety of suitable means: transformation, transfection, conjugation, protoplast fusion, electroporation, calcium phosphate-precipitation, direct microinjection, etc.

Host cells may be either prokaryotic or eukaryotic. Preferred are eukaryotic hosts, e.g. mammalian cells, such as human, monkey, mouse, and Chinese hamster ovary (CHO) cells, because they provide post-translational modifications to protein molecules, including correct folding or glycosylation at correct sites. Also yeast cells can carry out post-translational peptide modifications including glycosylation. A number of recombinant DNA strategies exist which utilize strong promoter sequences and high

10

15

20

25

-5-

copy number of plasmids which can be utilized for production of the desired proteins in yeast. Yeast recognizes leader sequences on cloned mammalian gene products and secretes peptides bearing leader sequences (i.e., pre-peptides).

After the introduction of the vector(s), the host cells are grown in a selective medium, which selects for the growth of vector-containing cells. Expression of the cloned gene sequence(s) results in the production of the desired proteins

The amino-terminal truncated chemokines of the invention may be prepared by any other well known procedure in the art, in particular, by the well established chemical synthesis procedures, utilizing automated solid-phase peptide synthesizers followed by chromatographic purification.

The chemokines of the invention may, for example, be synthesized by Fmoc (9-fluorenylmethoxycarbonyl), tBoc (t-butoxycarbonyl) or any other comparable chemical synthesis with or without appropriate side-chain protection groups on the different amino acids. The amino acids with or without appropriate side-chain protection groups are preactivated - e.g. with HBTU/HOBt [2-(1H-Benzotriazole-1yl)-1,1,3,3-tetramethyl-uromium hexafluorophosphate/1-hydroxybenzotriazole) - and coupled to the growing peptide chain. Before the addition of the following residue, the protection group (e.g. Fmoc) is removed from the α-amino group. After synthesis, all protection groups are removed, the intact full length peptides are purified and chemically or enzymatically folded (including the formation of disulphide bridges between cysteines) into the corresponding chemokines of the invention.

Purification of the natural, synthetic or recombinant proteins is carried out by any one of the methods known for this purpose, i.e. any conventional procedure involving extraction, precipitation, chromatography, electrophoresis, or the like (see for example Proost P. et al., Methods: A companion to Methods in Enzymol., 10, 82, 1996). A further purification procedure that may be used in preference for purifying the protein of the invention is affinity chromatography using monoclonal antibodies, affinity or heparin, which bind the target protein and which are produced and immobilized on a gel matrix contained within a column. Impute preparations containing the proteins are passed

10

15

20

25

-6-

while the impurities will pass through. After washing, the protein is eluted from the gel by a change in pH or ionic strength.

The amino-terminal truncated chemokines of the invention are useful in the therapy and/or diagnosis of the diseases, in which an antagonistic activity of the chemokine effects is required. Examples of such diseases include: inflammatory diseases, angiogenesis- and hematopoiesis-related diseases, tumors, infectious diseases, including HIV, auto-immune diseases, atherosclerosis, pulmonary diseases and skin disorders.

Therefore, in a further aspect, the present invention provides the use of the protein of the invention in the manufacture of a medicament for the treatment of the above-mentioned diseases.

The medicament is preferably presented in the form of a pharmaceutical composition comprising the proteins of the invention together with one or more pharmaceutically acceptable carriers and/or excipients. Such pharmaceutical compositions form yet a further aspect of the present invention.

It has also been found that CD26/DPP IV is able to generate NH<sub>2</sub>-terminally truncated chemokines in vitro. RANTES is the first cytokine reported whose biological activity can be modified by CD26/DPP IV.

Therefore, another object of the present invention is the use of CD26/DPP IV in the therapy and/or diagnosis of the diseases, in which an antagonistic activity of the chemokine effects is required, with particular focus on inflammatory, immune and infectious diseases.

Since this represents the first example of an identified mechanism for endogeneously regulated chemokine modification into an antagonist, similar physiological processing, but mediated by other factors (proteases), must be responsible for the MCP-2 truncation into MCP-2(6-76). The use of such factors (proteases) in the therapy and/or diagnosis of the above diseases is also included in this invention.

The invention will now be described by means of the following Examples, which should not be construed as in any way limiting the present invention. The Examples will refer to the Figures specified here below.

30

6

#### DESCRIPTION OF THE FIGURES

-7-

Figure 1: it shows the amino acid sequences of MCP-2 and RANTES. Signal sequences are reported in *italics*, whereas C -residues are in **bold**. Arrows indicate the first amino acids of the amino-terminal truncated chemokines of the invention.

Figure 2: SDS-PAGE of amino-terminal truncated MCP-2(6-76):

5 lane 1: natural MCP-2 (1-76, 100 ng/ml);

lane 2: natural MCP-2 (1-76, 30 ng/ml);

lane 3; natural MCP-2 (6-76, 30 ng/ml); and

lane 4: synthetic MCP-2 (1-76, 60 ng/ml).

Gels were run under reducing conditions and proteins were stained with silver.

- Figure 3: it shows a comparison of the chemotactic potency of modified MCP-2 forms. Intact natural and synthetic MCP-2(1-76), NH<sub>2</sub>-terminally truncated natural MCP-2(6-76) and COOH-terminally truncated synthetic MCP-2(1-74) were tested for chemotactic activity on THP-1 cells. Results represent the mean CI ± SEM from four or more independent experiments.
- Figure 4: Natural MCP-2 is a weaker agonist than MCP-1 to mobilize calcium in monocytes. Intact MCP-2 (15, 50 and 150 ng/ml) dose-dependently increases the [Ca<sup>2+</sup>]<sub>i</sub> in THP-1 cells. The result of one representative experiment out of two is shown.
- Figure 5: The chemotactic potencies of intact and NH<sub>2</sub>-terminally truncated forms of natural or recombinant RANTES for monocytic THP-1 cells were compared in the Boyden microchamber assay: Natural RANTES(1-68) (Δ), natural, truncated RANTES(3-68) (□), intact recombinant RANTES(1-68) (Δ) and CD26/DPP IV cleaved recombinant RANTES(3-68) (□). Results represent the mean chemotactic index ± SEM of four or more independent experiments.
- Figure 6: Effect of natural RANTES(3-68) (□), natural RANTES(1-68) (Δ), recombinant

  25 RANTES(1-68) (Δ) and recombinant CD26/DPP IV treated RANTES(3-68) (□) on the

  [Ca²¹], in THP-1 cells. Results represent the mean increase in [Ca²¹], ± SEM of three or more independent experiments.
  - Figure 7: It shows the desensitization of the Ca<sup>2+</sup>-mobilizing activity of intact

15

20

25

-8-

represent the mean  $\pm$  SEM (three or more independent experiments) increase in [Ca<sup>2+</sup>], in response to 30 ng/ml of intact recombinant RANTES as a second stimulus.

#### **EXAMPLES**

# 5 EXAMPLE 1: Amino-terminally truncated MCP-2

Materials and methods

Chemokine and immunoassay

MCP-2 was synthesized and purified as described earlier (Proost P. et al., Cytokine 7(97), 1995).

Specific anti-human MCP-2 Ab were obtained from mice and affinity purified on a Sepharose column to which synthetic MCP-2 was coupled using the conditions provided by the manufacturer (CNBr activated Sepharose 4B, Pharmacia, Uppsala, Sweden).

ELISA plates were coated with the affinity purified anti-human MCP-2 and biotinylated anti-MCP-2 was used as the capturing Ab. The detection was performed with peroxidase labeled streptavidine and TMB. The detection limit for MCP ELISA was about 0.1 ng/ml.

#### Production and purification of MCP-2

Monocyte chemotactic proteins were purified from peripheral blood mononuclear cell-derived conditioned medium from 132 blood donations obtained from Blood Transfusion Centers of Antwerp and Leuven (Proost P. et al., Methods: A companion to Methods in Enzymol., 10, 82, 1996).

Erythrocytes and granulocytes were removed by sedimentation in hydroxyethyl starch (Fresenius AG, Bad Homburg, Germany) and by gradient centrifugation in a sodium metrizoate solution (Lymphoprep; Nyegaard, Oslo Norway).

Mononuclear cells ( $60x10^9$  cells) were incubated ( $5x10^6$  cells/ml) with  $10 \mu g/ml$  Con A and 2  $\mu g/ml$  of LPS. After 48 to 120 h, conditioned medium was collected and kept at -20 °C until purification.

\_Q

Natural MCP was purified in a four step purification procedure as previously described (Proost P. et al., Methods: A companion to Methods in Enzymol., 10, 82, 1996).

Briefly, the conditioned medium was concentrated on controlled pore glass or silicic acid and partially purified by affinity chromatography on a heparin-Sepharose column (Pharmacia).

Fractions containing MCP immunoreactivity were further purified by Mono S (Pharmacia) cation exchange chromatography and eluted in a NaCl gradient at pH 4.0.

Natural MCPs were purified to homogeneity through RP-HPLC on a C-8

Aquapore RP-300 column (Perkin Elmer, Norwalk CT) equilibrated with 0.1 % trifluoroacetic acid TFA). Proteins were eluted in an acetonitrile gradient.

10

15

20

25

30

-1()-

Biochemical characterization of MCP-forms by SDS-PAGE, amino acid sequence analysis and mass spectrometry

The purity of column fractions was examined by SDS-PAGE under reducing conditions on Tris/tricine gels (Proost P. et al., Methods: A companion to Methods in Enzymol., 10, 82, 1996). Proteins were stained with silver and the following relative molecular (Mr) markers were used: OVA (Mr 45,000), carbonic anhydrase (Mr 31,000), soybean trypsin inhibitor (Mr 21,500), β-lactoglobulin (Mr 18,400), lysozyme (Mr 14,400) and aprotinin (Mr 6,500).

The NH<sub>2</sub>-terminal sequence of purified chemokines was determined by Edman degradation on a pulsed liquid 477A/120A protein sequencer (Perkin Elmer) with N-methylpiperidine as a coupling base. Blocked proteins were cleaved between Asp and Pro in 75 % formic acid for 50 h. The formic acid digest was sequenced without further purification.

The Mr of MCP-2 was determined by matrix-assisted laser desorption ionization/time of flight-mass spectrometry (MALDI/TOF-MS) (Micromass TofSpec, Manchester, UK). Alpha-cyano-4-hydroxycinnamic acid and cytochrome C were used as matrix and internal standard, respectively.

#### Detection of chemotactic activity

MCP-2 was tested for its chemotactic potency on freshly purified monocytes  $(2\times10^6 \text{ cells/ml})$  or monocytic THP-I cells  $(0.5\times10^6 \text{ cells/ml})$ ; 2 days after subcultivation) in the Boyden microchamber using polyvinylpyrrolidone-treated polycarbonate membranes with 5  $\mu$ m pore size.

Samples and cells were diluted in HBSS (Life technologies/Gibco BRL, Paisley, Scotland) supplemented with 1 mg/ml human serum albumin (Red Cross Belgium). After 2 h incubation at 37 °C, the cells were fixed and stained with Diff-Quick staining solutions (Harleco, Gibbstown. NJ) and the cells that migrated through the membranes were counted microscopically in ten oil immersion fields at 500-fold magnification.

The chemotactic index (CI) of a sample (triplicates in each chamber) was calculated as the number of cells that migrated to the sample over the number of cells that migrated to control medium (Van Damme J. J. Exp. Med., 176, 59, 1992).

15

20

2.5

-11-

For desensitization experiments, cells were incubated with biologically inactive chemokine-variants for 10 min. at 37°C, before they were added to the upper well of the Boyden microchamber. The % inhibition of the CI was calculated using the CI from HBSS treated cells towards the sample as a reference value.

Detection of intracellular Ca2+ concentrations

Intracellular calcium concentrations ([Ca<sup>2+</sup>]<sub>i</sub>) were measured as previously described (Wuyts et al., *Biochemistry*, 32, 2716, 1997). Purified monocytes or THP-1 cells (107 cells/ml) were incubated in Eagle's Minimum Essential Medium (EMEM, Gibco) + 0.5 % FCS with the fluorescent indicator fura-2 (fura-2/AM 2.5 µM; Molecular Probes Europe BV, Leiden, The Netherlands) and 0.01 % Pluronic F-127 (Sigma, St Louis MO).

After 30 min at 37 °C the cells were washed twice and resuspended at 106 cells/ml in HBSS with 1 mM Ca<sup>2+</sup> and 0.1 % FCS (buffered with 10 mM Hepes/NaOH at pH 7.4). The cells were equilibrated at 37 °C for 10 min before fura-2 fluorescence was measured in a LS50B luminescence spectrophotometer (Perkin Elmer).

Upon excitation at 340 and 380 nm, fluorescence was detected at 510 nm. The  $[Ca^{2^{1}}]_{i}$  was calculated from the Grynkiewicz equation (Grynkiewicz et al., *J. Biol. Chem.*, 260, 3440, 1985). In order to determine  $R_{max}$  the cells were lysed with 50  $\mu$ M digitonin. Subsequently, the pH was adjusted to 8.5 with 20 mM Tris and  $R_{min}$  was obtained by addition of 10 mM EGTA to the lysed cells. The  $K_{d}$  used was 224 nM.

For desensitization experiments, monocytes or THP-1 cells were first stimulated with buffer, chemokine or chemokine antagonist at different concentrations. As second stimulus, MCP-2 was used at a concentration inducing a significant increase in the [Ca<sup>2+</sup>], after prestimulation with buffer. The second stimulus was applied 2 min after addition of the first stimulus. The percentage inhibition of the [Ca<sup>2+</sup>]; increase in response to the second stimulus was calculated comparing the signal after prestimulation with chemokine or chemokine antagonist with the signal after addition of buffer.

10

15

20

25

30

-12-

Results

Isolation of post-translationally modified MCP-2 forms

A specific and sensitive ELISA was used to trace different MCP-2 forms produced by peripheral blood mononuclear cells stimulated with mitogen and endotoxin.

The conditioned medium was purified according to a standard isolation procedure (Proost P. et al., Methods: A companion to Methods in Enzymol., 10, 82, 1996), including adsorption to controlled pore glass and heparin Sepharose chromatography.

Subsequently, purification by FPLC mono S cation exchange chromatography was carried out and then a further purification step with C-8 RP HPLC was applied. Molecular masses were measured by SDS-PAGE and by MALDI/TOF-MS.

Different forms of MCP-2 were isolated, in addition to the authentic 7.5 kDa MCP-2(1-76), an NH<sub>2</sub>-terminally truncated 7 kDa form of MCP-2 missing five residues [MCP-2(6-76)] was purified to homogeneity by RP-HPLC and identified by amino acid sequence analysis (Fig. 2). MALDI/TOF-MS (Table I) yielded a molecular mass of 8881 Da for intact MCP-2 (theoretical *Mr* of 8893 Da), whereas for the MCP-2(6-76) a molecular mass of 8365 Da was measured, confirming the deletion of the five NH<sub>2</sub>-terminal amino acids (theoretical *Mr* of 8384 Da). Functional comparison of these natural MCP-2 forms in the THP-1 chemotaxis assay showed that intact MCP 2 is still active at 5 ng/ml, whereas truncated MCP-2(6-76) remains devoid of chemotactic activity when tested at a concentration range from 0.6 to 60 ng/ml (Fig. 3). Intact natural MCP-2 was also compared in potency with the synthetical MCP-2(1-76) and a COOH-terminally truncated synthetical form (Proost P. et al., *Cytokine* 7(97), 1995) missing two residues [MCP-2(1-74)].

The minimal effective chemotactic concentration of these forms was also found to be 5 ng/ml (Fig. 3). Although in chemotaxis assays the specific activity of natural intact MCP-1 and MCP-2 is comparable (Van Damme J, et al., *J. Exp. Med.*, 176, 59, 1992), the calcium mobilizing capacity of MCP-2 is still a matter of debate.

However, in Ca<sup>2+</sup> -mobilization experiments, the minimal effective dose for both natural or synthetic MCP-2(1-76) was 10-fold higher compared to that of natural intact MCP-1(1-76) (Fig. 4), whereas MCP-2(6-76) remained inactive.

10

15

-13-

Nevertheless, intact MCP-2 (50 ng/ml) was capable to desensitize for MCP-2 (15 ng/ml) and MCP-3 (10 ng/ml) yielding 52% and 45% inhibition, respectively.

Due to this lower specific activity of MCP-2 in Ca<sup>2+</sup> assays, desensitization of chemotaxis by MCP-2(6-76) was performed in the Boyden microchamber. Since intact MCP-2 is reported to cross-desensitize with active MCP-1, MCP-2 and MCP 3 in the monocyte chemotaxis assay (Sozzani et al., *J. Immunol.*, 152, 3615, 1994), we investigated whether natural, inactive MCP-2(6-76) could also desensitize for MCP-1, MCP-2, MCP-3 and RANTES (Table III). Pre-incubation of THP-I cells with 100 ng/ml of inactive MCP-2(6-76) could already significantly inhibit chemotaxis induced by 10 ng/ml of MCP-1 (63 %), 5 ng/ml of MCP-2 (75%), 30 ng/ml of MCP-3 (62 %) and 100 ng/ml of RANTES (75%). Moreover, chemotaxis in response to 3 times lower concentrations of the respective MCPs was completely (91-100 %) inhibited by 100 ng/ml MCP-2(6-76). Furthermore, at a concentration as low as 10 ng/ml, MCP-2(6-76) was still able to significantly inhibit the chemotactic activity induced by MCP-1 (3 ng/ml), MCP-2 (1.5 ng/ml) or MCP 3 (10 ng/ml) and RANTES (30 ng/ml). Taken together, MCP-2(6-76) is produced naturally, is inactive as a chemoattractant and antagonizes several C-C chemokines, the effect being most predominant for MCP-3.

#### TABLE I

20

Biochemical characterization of natural forms of MCP-2. NH2-terminal amino acid sequence analysis and comparison of the experimental (SDS-PAGE and MALDI/TOF-MS) and theoretical Mr of C-8 RP-HPLC purified natural MCP-isoforms.

MCP-form	NH <sub>2</sub> -terminal sequence	Mr (Da)		
		theoretical unglycosylated	SDS-PAGE	MALDI/TOF-MS
MCP-2 (1-76)	blocked	8893	7500	8881

-14-TABLE II

MCP-2(6-76) desensitizes the monocyte chemotactic responses of MCP-1, MCP-2 MCP-3 and RANTES in the microchamber.

Chemokine "	Concentration	Antagonizatio	% Inhibition of	
		resp	ons <b>e</b> <sup>A.c</sup>	chemotaxis
		buffer	100 ng/ml	
			MCP-2(6-76)	
MCP-1	10	22.3 ± 7.9	8.3 ± 3.8	63 ± 21
	3	15.0 ± 8.0	$1.3 \pm 0.3$	99 ± 1.0
MCP-2	5	$36.0 \pm 12.6$	10.8 ± 6.1	75 ± 8.0
	1.5	6.7 ± 1.4	$1.5 \pm 0.3$	91 ± 70
MCP-3	30	13 2 ± 0.4	6.0 ± 4.0	62 ± 31
	10	$3.0 \pm 1.5$	<1	100 ± 0.0
RANTES	100	6.3 ± 0.8	2.6 ± 1.3	75 ± 19
	30	$4.0 \pm 0.8$	$1.5 \pm 0.3$	77 ± 1.6
		buffer	10 ng/ml	
			MCP-2 (6-76)	
MCP-1	10	12.7 ± 2.3	$10.5 \pm 3.8$	24 ± 1.8
	3	$7.5 \pm 0.0$	$3.0 \pm 0.3$	69 ± 4.0
MCP-2	5	$38.0 \pm 5.3$	$27.2 \pm 4.9$	$30 \pm 6.0$
	1.5	18.3 ± 4.6	9.2 ± 1.4	45 ± 23
MCP-3	30	$13.2 \pm 1.9$	$8.0 \pm 1.0$	37 ± 19
	10	7.7 ± 1.4	$1.7 \pm 0.3$	90 ± 6.0
RANTES	100	5.5 ± 0.6	$5.8 \pm 0.9$	17 ± 7.0
	30	$3.2 \pm 0.7$	$2.5 \pm 0.5$	39 ± 18

MCP-1, MCP-2, MCP-3 or RANTES were added as chemoattractants to the lower wells.

b the upper wells of the microchamber were filled with THP-1 cells preincubated with MCP-2(6-76) or with buffer

<sup>\*</sup> mean CI ± SEM of 3 independent experiments

15

20

25

-15-

## **EXAMPLE 2: Amino-terminally truncated RANTES**

Reagents

Natural human RANTES was produced by human Malavu hepatosarcoma cells, MG-63 osteosarcoma cells or peripheral blood leukocytes (Blood transfusion centers of Antwerp and Leuven) and purified as previously described (Proost, P. et al., (1996) Methods: A Companion to Methods in Enzymol. 10, 82-92; and Proost, P. et al., (1993) J. Immunol. 150, 1000-1010). MCP-2, MCP-3 and GCP-2 were synthesized by Fmoc chemistry (Proost, P. et al., (1995) Cytokine 7, 97-104; and Wuyts, A. et al., (1997) Biochemistry 36, 2716-2723), recombinant human RANTES was obtained from Peprotech (Rocky Hill, NJ) and recombinant MCP-1 was a gift from Dr. J.J. Oppenheim (NCI-NIH, Frederick, MD).

Human CD26/DPP IV was obtained from prostasomes, prostate derived organelles, which occur freely in seminal plasma. The enzyme was purified to homogeneity as described before using ion exchange followed by affinity chromatography onto adenosine deaminase (De Meester, I. et al., (1996) J. Immunol. Methods 189, 99-10526).

Incubation of chemokines with CD26/DPP IV and detection of proteolytic processing

A 100 to 1000 molar excess of chemokine was incubated overnight with CD26/DPP IV in 100 mM Tris/HCl pH 7.7. Chemokines were separated from CD26/DPP IV by SDS-PAGE on a Tris/Tricine gel system as previously described (Proost, P. et al., (1996) Methods: A Companion to Methods in Enzymol. 10, 82-92).

Proteins were electroblotted on PVDF (polyvinylidene fluoride) membranes (Problott, Perkin Elmer, Foster City, CA) and stained with coomassie brilliant blue R250. After destaining, membranes were rinsed at least 5 times with ultrapure water (Milli Q; Millipore, Bedford, MA).

To obtain sufficient amounts of pure truncated chemokine for biological assays, about 50 µg of recombinant chemokine was treated with CD26/DPP IV and the cleavage product was acidified with 0.1.% trifluoroacetic acid (TFA). Tween 20.00.1.%) was

separated from CD26-DPP IN the an acetomitrile gradient to be the Aquapore RP-Not

10

15

20

30

-16-

column (1 x 50 mm) (Perkin Elmer). Fractions containing proteins were analyzed by SDS-PAGE and silver stained as described.

CD26/DPP IV treated chemokines, purified by RP-HPLC or excised from PVDF blots, were NH<sub>2</sub>-terminally sequenced by Edman degradation on a pulsed liquid phase 477A/120A protein sequencer (Perkin Elmer) using N-methylpiperidine as a coupling base.

#### Detection of chemotactic activity

Chemokines were tested for their chemotactic potency on freshly isolated peripheral blood neutrophilic granulocytes (10<sup>6</sup> cells/ml) or cultured monocytic THP-1 cells (0.5 x 10<sup>6</sup> cells/ml) in the Boyden microchamber (Proost, P. et al., (1996) *Methods:* A Companion to Methods in Enzymol. 10, 82-92; and Proost, P. et al., (1993) J. Immunol. 150, 1000-1010).

After 45 min (granulocytes) or 2 h (THP-1 cells) incubation at 37 °C, the cells were fixed and stained. The cells that migrated through the 5 µm pore size polycarbonate membranes were counted microscopically in ten oil immersion fields. The chemotactic index (C.I.) of a sample (triplicates in each chamber) was calculated as the number of cells that migrated to the test sample divided by the number of cells that migrated to control medium. In desensitization experiments, cells were incubated with biologically inactive chemokine-variants for 10 min at 37 °C before transfer to the chamber.

The percentage inhibition of the C.I. obtained by desensitization with HBSS-treated control cells was calculated for the evaluation of chemotaxis desensitization.

# 25 Detection of intracellular Ca2+ concentrations

Intracellular Ca<sup>2+</sup> concentrations ([Ca<sup>2+</sup>]<sub>i</sub>) were measured as previously described (Wuyts, A., et al., (1997) *Biochemistry* 36, 2716-2723). Briefly, purified cells were incubated with the fluorescent indicator fura-2 (2.5 µM fura-2/AM, Molecular Probes Europe BV, Leiden, The Netherlands) and 0.01 % Pluronic F-127 (Sigma, St. Louis, MO).

10

15

20

25

and the region of the state of the same

-17-

After 30 min, cells were washed twice, resuspended in HBSS with 1 mM Ca<sup>2+</sup> and incubated for 10 min at 37 °C before fura-2 fluorescence was measured in an LS50B luminescence spectrophotometer (Perkin Elmer). Upon excitation at 340 and 380 nm, fluorescence was detected at 510 nm. The [Ca<sup>2+</sup>]<sub>i</sub> was calculated from the Grynkiewicz equation (Grynkiewicz, G. et al. (1985) J. Biol. Chem. 260, 3440-3450).

In order to determine  $R_{\rm max}$ , the cells were lysed with 50  $\mu$ M digitonin. Subsequently, the pH was adjusted to 8.5 with 20 mM Tris and  $R_{\rm min}$  was obtained by addition of 10 mM EGTA to the lysed cells. The  $K_a$  used for calibration was 224 nM. For desensitization experiments, cells were first stimulated with buffer or chemokine at different concentrations. As a second stimulus, chemokines were added at a concentration inducing a significant increase in the  $[Ca^{2-}]_i$  after prestimulation with buffer. The percentage inhibition of the  $[Ca^{2-}]_i$ -increase in response to the second stimulus by prestimulation of the cells was calculated.

#### Results

Identification and biological characterization of natural, NH<sub>2</sub>-terminally truncated RANTES.

A different NH<sub>2</sub>-terminally truncated form of human GCP-2 has been previously isolated (Proost, P. et al. (1993) J. Immunol. 150, 1000-1010). The least truncated GCP-2-form was cleaved beyond Pro at the penultimate position [GCP-2(3-75)]. Using a similar standard purification procedure (cfr. also MCP-2, vide supra), the C-C chemokine RANTES was purified from peripheral blood leukocytes or sarcoma cells (Proost, P. et al., (1996) Methods: A Companion to Methods in Enzymol. 10, 82-92).

In addition to full length human RANTES, a truncated RANTES-variant missing the two NH<sub>2</sub>-terminal residues [RANTES(3-68)], was consistently isolated.

RANTES(3-68) was tested for chemotactic and/or intracellular Ca<sup>2+</sup>-releasing activity and their biological potency was compared with that of the respective intact chemokines.

haturai RANTES (minimai effective dose of 5% ng/m), naturai RANTES(3/08) was

15

20

25

30

-18-

totally inactive when tested at concentrations as high as 300 ng/ml in the Boyden microchamber (Fig. 5). In addition, 10 times higher concentrations of natural RANTES(3-68), compared to RANTES(1-68), were necessary to obtain a similar Ca<sup>2+</sup>-response (Fig. 6).

5 CD26/DPP IV removes the NH<sub>2</sub>-terminal dipeptides of chemokines

In order to investigate whether the aminopeptidase CD26/DPP IV could be responsible for the NH<sub>2</sub>-terminal truncation of RANTES, the intact chemokine was incubated overnight with CD26/DPP IV, blotted to PVDF membranes, stained with Coomassie blue and subjected to automatic Edman degradation. CD26/DPP IV treatment of RANTES resulted in the removal of the NH<sub>2</sub>-terminal dipeptides. Parallel incubation of chemokine with buffer without CD26/DPP IV had no effect.

Since other chemokines contained the consensus sequence for CD26/DPP IV cleavage and since the NH<sub>2</sub>-terminus of MCPs was shown to be crucial for biological activity (Gong, J. et al. (1996) J. Biol. Chem. 271, 10521-10527, and Gong, J. et al. (1995) J. Exp. Med. 181, 631-640), MCP-1, MCP-2 and MCP-3 were also incubated with CD26/DPP IV.

After treatment, MCPs were blotted on PVDF membranes and Coomassie blue stained to confirm that a sufficient amount of protein was recovered for Edman degradation.

However, no NH<sub>2</sub>-terminal sequence could be detected, indicating that CD26/DPP IV does not alter the NH<sub>2</sub>-terminus of MCPs which is blocked for Edman degradation by a pyroglutamic acid.

Comparison of the biological activity of intact and CD26/DPP IV-treated RANTES.

Similar to natural RANTES(3-68), C-8 RP-HPLC purified, CD26/DPP IV-treated recombinant RANTES was inactive in Boyden microchamber chemotaxis experiments when used at concentrations up to 1 µg/ml, while a significant monocyte chemotactic response was detected with intact recombinant RANTES from 30 to 100 ng/ml onwards (Fig. 5).

When the truncation effect was tested in the Ca<sup>2-</sup>-mobilization assay, RANTES(3-68) induced a low but significant increase at 100 ng/ml. Intact RANTES,

15

-19-

however, was already active at 10 ng/ml (Fig. 6). In conclusion, although only two NH<sub>2</sub>-terminal residues were removed, the monocyte chemotactic and Ca<sup>2</sup>'-mobilizing potency of RANTES decreased 10 to 100-fold.

## 5 RANTES(3-68) is a natural chemotaxis antagonist for intact RANTES

In view of the inactivity of RANTES(3-68) in monocyte chemotaxis experiments, we tested whether this truncated RANTES might act as an antagonist. RANTES(3-68), at 1 µg/ml, almost completely (82 %) desensitized for the chemotactic effect of 100 ng/ml of intact RANTES (Table III).

When a 3-fold excess of RANTES(3-68) was added to the upper well, chemotaxis of THP-1 cells towards intact RANTES was inhibited by about 50-70 %. RANTES(3-68) at 300 ng/ml could still inhibit about 30 % of the chemotactic response towards an equal concentration of intact RANTES.

In  $Ca^{2+}$ -mobilization experiments with THP-1 cells (Fig. 7), 30 ng/ml of intact RANTES could desensitize for the effect of 30 ng/ml of intact RANTES for 39  $\pm 5$  %. About ten-fold higher concentrations of RANTES(3-68) were necessary to obtain the same amount of desensitization. However, at 300 ng/ml, RANTES(3-68) by itself gave a significant  $Ca^{2+}$ -response. This  $Ca^{2+}$ -response was comparable to the response obtained with 30 ng/ml of intact RANTES.

-20-TABLE III

RANTES(3-68) desensitizes monocyte chemotaxis induced by RANTES(1-68)1

Chemokine(ng/ml)		Chemotactic response (CI)					
Lower well	Upper well			% inhibition			
RANTES (1-68)	RANTES (3-68)	A	B	С	D	mean ± SEM	mean ± SEM
300	1000	12.5	7.5	27.5	50.5	25 ± 10	67 ± 8
	300	22.0	20,5	72.5	79.5	49 ± 16	31 ±13
	0	41.0	46.0	71.5	97.0	64 ±1 3	υ
100	1000	4.0	3.0	13.5	11.0	8 ± 3	82 ± 4
	300	7.5	7.0	29.0	33.0	19±7	53 ± 11
	0	24.0	21.5	50.0	44.5	35 ± 7	0

<sup>&</sup>lt;sup>1</sup> Results represent the chemotactic index (C.I.) of four (A to D) independent experiments (including mean ± SEM) and the percentage (%) inhibition (mean ± SEM of the % inhibition of the four experiments) of the chemotactic response towards

10 RANTES(1-68) after preincubation of the THP-1 cells with inactive RANTES(3-68) or buffer.



# -21-CLAIMS

- 1. Amino-terminal truncated C-C chemokines, having chemokine antagonistic activity.
- 2. Amino-terminal truncated C-C chemokines, according to claim 1, missing up to 5 amino-terminal amino acids and having a chemokine antagonistic activity
  - 3. Amino-terminal truncated C-C chemokines, according to any of the preceding claims selected among the group consisting of MCP-1, MCP-2, MCP-3, MCP-4, MCP-5 and RANTES.

10

5

- 4. Amino-terminal truncated MCP-2(6-76), which is MCP-2 missing the 5 NH<sub>2</sub>-terminal amino acids, as shown in Figure 1.
- 5. Amino-terminal truncated RANTES(3-68), which is RANTES missing the first 2 amino acids, as shown in Figure 1.
  - 6. Amino-terminal truncated C-C chemokines, according to one or more of the preceding claims, in a glycosylated form.
- 7. DNA molecules comprising the DNA sequences coding for the amino-terminal truncated chemokines of the invention according to one or more of the preceding claims, including nucleotide sequences substantially the same
  - 8. An expression vector which comprises the DNA molecule of any claim 7.
- 25 9. A host cell comprising the expression vector of claim 8.
  - 10.A recombinant process for preparing any of the proteins from claim 1 to 5, comprising culturing in an appropriate culture medium the cells of claim 9.

15

-22-

- 12. Use of a protein according to any of the claims from 1 to 6, in the manufacture of a medicament for the therapy and/or diagnosis of diseases, in which an antagonistic activity of the chemokine effects is required.
- 5 13.Use according to claim 12, in the manufacture of a medicament for the treatment of inflammatory diseases, HIV-infection, angiogenisis- and hematopoiesis-related diseases, and tumors.
  - 14.A pharmaceutical composition comprising the protein according to any of the claims from 1 to 6 together with one or more pharmaceutically acceptable carriers and/or excipients.
  - 15.Use of CD26/DPP IV in the therapy and/or diagnosis of the diseases, in which an antagonistic activity of the chemokine effects is required.
  - 16.Use according to claim 15, for the treatment of inflammatory, immune and infectious diseases.

-23-

#### **ABSTRACT**

Amino-terminal truncated C-C chemokines, having chemokine antagonistic activity, are described. In particular, we refer to amino-terminal truncated C-C chemokines missing up to 5 amino-terminal amino acids and having a chemokine antagonistic activity. More particularly, MCP-2 (6-76) and RANTES (3-68) are described. The cDNA sequences encoding them, their use in therapy and/or in diagnosis of the diseases, in which an antagonistic activity of the chemokine effects is required, as well as pharmaceutical compositions comprising them.





RANTES

-23

1 ↓

1

MKVSAARLAV ILIATALCAF ASASPYSSDT TPCCFAYIAR PLPRAHIKEY PYTSGKCSNP AVVFVTRKNR QVCANPEKKW VREYINSLEM S

58

1

MCP2

-23

MKVSAALLCL LLMAATFSPQ GLAQPDSVSI PITCCFNVIN RKIPIQRLES YTRITNIQCP

KEAVIFKTKR GKEVCADFKE RWVRDSMKHL DQIFQNLKF

76

Figure 1

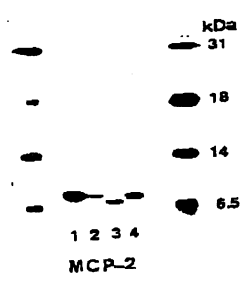


Figure 2

SPEC

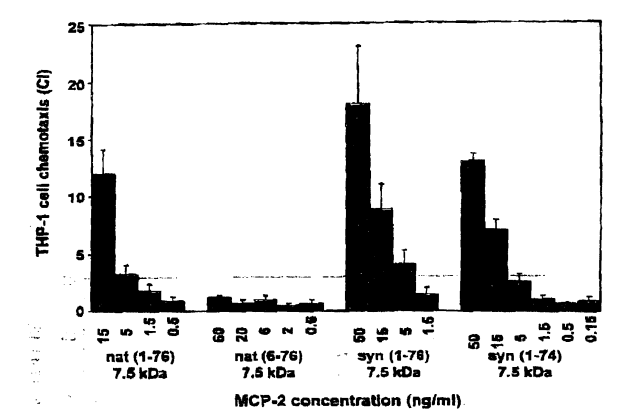
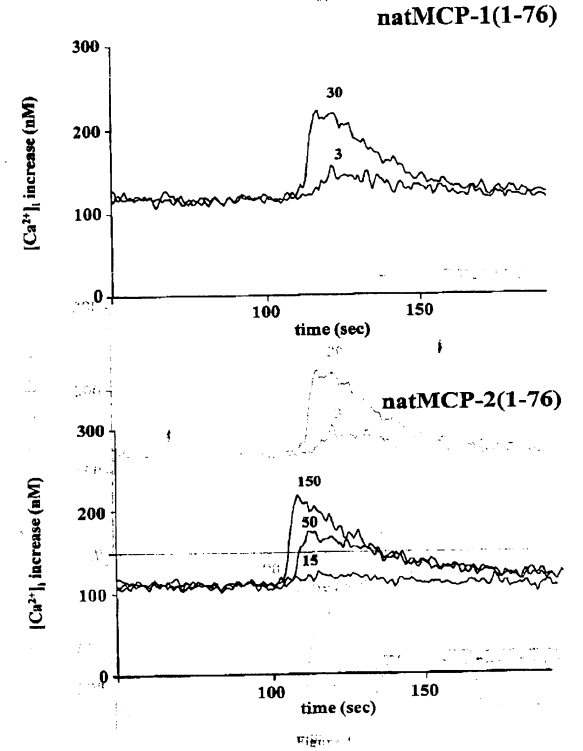


Figure 3





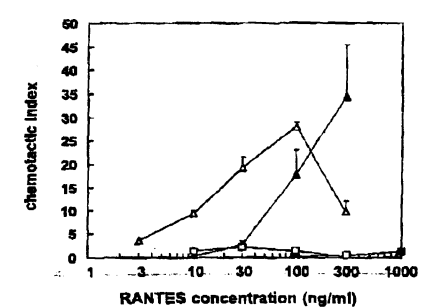


Figure 5

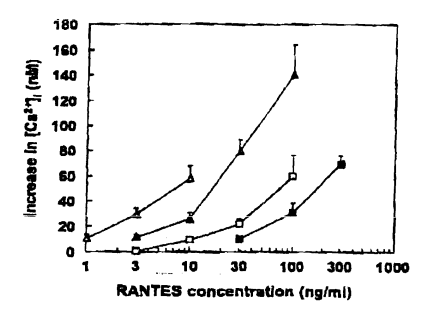


Figure 6

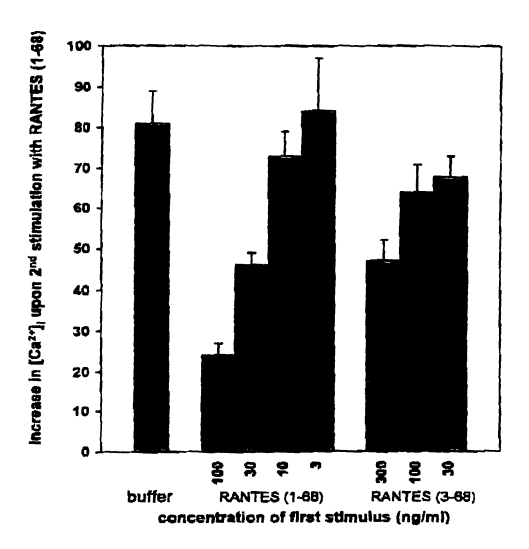


Figure 7